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(1) Publication number:

<mark>0 295 627</mark> A1

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EUROPEAN PATENT APPLICATION

- 2 Application number: 88109469.2
- 1 Int. Cl.4 A61K 6/08

- Date of filing: 14.06.88
- Priority: 19.06.87 JP 163022/87
- ② Date of publication of application: 21.12.88 Bulletin 88/51
- Designated Contracting States:
 DE FR GB

- Applicant: MITSUBISHI RAYON CO. LTD. 3-19, Kyobashi 2-chome Chuo-Ku Tokyo 104(JP)
- (7) Inventor: Sasaki, Isao
 1665-98, Ohno-cho
 Saiki-gun Hiroshima-ken(JP)
 Inventor: Mukai, Nobuhiro
 13-19-105, Inokuchidai 1-chome
 Nishi-ku Hiroshima-shi Hiroshima-ken(JP)
 Inventor: Ige, Hitoshi
 3-16-303, Nishisakae 3-chome
 Ohtake-shi Hiroshima-ken(JP)
- Representative: Patentanwäite TER MEER MÜLLER STEINMEISTER
 Mauerkircherstrasse 45
 D-8000 München 80(DE)
- Photopolymerizable dental composition.
- A photopolymerizable dental composition, which comprises:
 (A) an organic composite filler obtained by polymerizing at least one carboxylic acid monomer of the formula:

$$R_2 = C = C$$

$$C = C$$

$$COOX$$
(I)

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wherein R₁ is a hydrogen atom, a C₁-C₁₅ alkyl group, a -COOY group, a halogen atom, or a phenyl group or a derivative thereof; R₂ is a hydrogen atom, a C₁-C₁₅ alkyl group, a -COOZ group, a halogen atom, or a phenyl group or a derivative thereof; R₃ is a hydrogen atom, a C₁-C₁₅ alkyl group, a halogen atom, or a phenyl group or a derivative thereof; and each of X, Y and Z is a hydrogen atom, NH₄ or an alkali metal atom; or the formula:

wherein each of R₄ and R₅ is a hydrogen atom, a C₁-C₁₅ alkyl group, a halogen atom, or a phenyl group or a derivative thereof; and at least one radical polymerizable vinyl monomer in a polymerization system in which an inorganic compound is dispersed;

(B) a monomer mixture comprising 2,2-bis(4-(methacryloxyethoxy)phenyl]propane, at least one hexafunctional urethane (meth)acrylate of the formula:

wherein X is
$$-(CH_2)$$
 HNCOO-CH $-(CH_2)$ CH2OOC-C= $-(CH_2)$ CH2OOC-C= $-(CH_2)$

wherein n is an integer of from 1 to 10, and each of R₁ and R₂ which may be the same or different is a hydrogen atom or a methyl group, and a diluting ethylenic vinyl monomer; and

(C) a photopolymerization initiator comprising an α -diketone compound and a reducing agent, wherein the reducing agent is a diaminobenzophenone compound of the formula:

$$R_1 R_2 N \longrightarrow C \longrightarrow NR_3 R_4$$
 (IV)

wherein each of R_1 to R_2 which may be the same or different is a hydrogen atom or a C_1 - C_2 alkyl group, provided that at least one of them is the alkyl group.

PHOTOPOLYMERIZABLE DENTAL COMPOSITION

The present invention relates to a dental filling composition containing a substantial amount of an organic composite filler as inorganic material. More particularly, it relates to a photopolymerizable dental filling composition which can readily be polymerized by inadiation with a visible light to provide a cured product having high mechanical properties and excellent water resistance.

For dental filling material, it is important that such material has not only good physical properties such as mechanical strength, abrasion resistance, water resistance and adhesive properties but also aesthetical properties such as transparency and polishability to make it look like natural teeth. Many attempts have been made to develop a so-called composite resin having an inorganic filler incorporated into a resin, as a substitute material for amalgam and with an object to improve the above-mentioned properties.

This composite resin is usually composed essentially of three components of (1) an inorganic filler, (2) an definic unsaturated compound and (3) a polymerization initiator. However, if an inorganic filler and an organic resin are simply blended, it is difficult to obtain adequate composite properties since the inorganic filler and the organic resin are poor in the interfacial affinity such as the compatibility or the adhesion to each other. In order to overcome such a drawback, a dental material has been proposed in which a filler prepared by treating a glass surface with a silane coupling agent is incorporated. However, the surface treatment with a silane coupling agent is limited in its applicable inorganic material to a glass filler having silanol groups on its surface and thus has a serious drawback that it is poor in the general applicability. Further, there is a problem such that the formed siloxane linkage is susceptible to hydrolysis and thus is poor in the water resistance. Accordingly, when applied to a dental filling composition, it hardly provides an interfacial reinforcing property to a quartz filler which is commonly employed as a filler to impart primarily high hardness and aesthetical properties. Further, it has a clinical problem such that since the water resistance is poor, the mechanical strength tends to deteriorate as time passes during clinic for a long period of time in a wet oral cavity.

Further, various studies have been made on the monomer constituting the organic resin. For example, a bisphenoi A multi-functional monomer represented by bisphenoi A diglycldyl methacrylate (hereinafter referred to simply as Bis-GMA) has been proposed as a monomer which provides a composite having a small polymerization shrinkage factor, and a hydrophobic monomer 2,2-bis[4-(methacryloxyethoxy)-phenyl]-propane (hereinafter referred to simply as Bis-MEPP) has been proposed as a monomer which provides excellent water resistance. However, Bis-GMA has a hydroxyl group in its molecule, and the resulting cured product is highly water absorptive and thus is poor in the water resistance. On the other hand, when Bis-MEPP is used, the curability is poor and adequate crosslinking density is hardly obtainable, and there has been a problem that the product is poor in the mechanical strength. Further, even if Bis-GMA and Bis-MEPP are combined to complement these drawbacks, it is impossible to obtain a composite resin having satisfactory properties. Accordingly, in order to determine a monomer composition for a particular purpose, it is necessary to study a very wide range of monomer compositions.

It is an object of the present invention to provide a photopolymerizable dental composition having excellent mechanical strength and good water resistance.

The present invention provides a photopolymerizable dental composition, which comprises:

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(A) an organic composite filler obtained by polymerizing at least one carboxylic acid monomer of the formula:

$$R_2 = C = C$$

$$C = C$$

$$Coox$$
(1)

wherein R₁ is a hydrogen atom, a C₁-C₁₅ alkyl group, a -COOY group, a halogen atom, or a phenyl group or a derivative thereof; R₂ is a hydrogen atom, a C₁-C₁₅ alkyl group, a -COOZ group, a halogen atom, or a phenyl group or a derivative thereof; R₂ is a hydrogen atom, a C₁-C₁₅ alkyl group, a halogen atom, or a phenyl group or a derivative thereof; and each of X, Y and Z is a hydrogen atom, NH₄ or an alkali metal atom; or the formula:

wherein each of R₄ and R₅ is a hydrogen atom, a C₁-C₁₅ alkyl group, a halogen atom, or a phenyl group or a derivative thereof; and at least one radical polymerizable vinyl monomer in a polymerization system in which an inorganic compound is dispersed;

(B) a monomer mixture comprising 2,2-bis[4-(methacryloxyethoxy)phenylipropane, at least one hexafunctional urethane (meth)acrylate of the formula:

$$\begin{array}{c|c}
x & C & X \\
\hline
 & C & X \\
\hline
 & C & X
\end{array}$$
(III)

wherein X is

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. 30

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$$: -(CH2) = HNCOO-CH CH2OOC-C=CH2 CH2OOC-C=CH2 R2$$

- wherein n is an integer of from 1 to 10, and each of R₁ and R₂ which may be the same or different is a hydrogen atom or a methyl group, and a diluting ethylenic vinyl monomer; and
 - (C) a photopolymerization initiator comprising an α-diketone compound and a reducing agent, wherein the reducing agent is a diaminobenzophenone compound of the formula:

$$R_1R_2N \longrightarrow C \longrightarrow NR_3R_4 \qquad (IV)$$

wherein each of R₁ to R₄ which may be the same or different is a hydrogen atom or a C₁-C₅ alkyl group, provided that at least one of them is the alkyl group.

Now, the present invention will be described in detail with reference to the preferred embodiments.

The organic composite filler (A) constituting the composition of the present invention can be obtained by polymerizing the carboxylic acid monomer with at least one other polymerizable vinyl monomer in a polymerization system in which an inorganic compound is dispersed. There is no particular restriction as to the method for its preparation. As a preferred embodiment, a method may be mentioned wherein a vinyl monomer and an inorganic compound are suspended and dispersed in an aqueous medium under such a

temperature condition that no thermal polymerization takes place, and then, the carboxylic acid monomer is added and stirred to initiate aqueous heterogeneous polymerization, whereby the polymerization is conducted for a predetermined period of time.

In the above polymerization, a radical polymerization initiator may be added as the case requires. For example, it is preferred to add a radical polymerization initiator when a high graft rate is required or when a mixture of various vinyl monomers is used.

The specific carboxylic acid monomer represented by the formula I or II to be used for the formation of the organic composite filler (A) is required to have a carboxylic acid group as an active site which brings about the polymerization activity and a double bond as an active site which provides firm consolidation of the resulting polymer and the inorganic compound. Any compound of the formula I or II having such functional groups can be used. Specifically, acrylic acid, methacrylic acid, crotonic acid, tiglic acid, cinnamic acid, maleic anhydride and citraconic anhydride may be mentioned. Among them, acrylic acid, methacrylic acid, crotonic acid and maleic anhydride are particularly preferred since they have very high polymerization activities.

The inorganic compound to be used for the preparation of the organic composite filler (A) includes elements of Groups I, II, III, IV and V of the Periodic Table, transition metals and their oxides, hydroxides, chlorides, sulfates, sulfites, carbonates, phosphates and silicates, and mixtures and complex salts thereof. Among them, barium sulfate, barium fluoride, silicon dioxide, alminum oxide, titanium oxide, quartz powder, glass powder, glass beads, glass fibers, a glass filler containing a barium salt, a lead salt or a strontium salt, silica gel, zirconium oxide and tin oxide are particularly preferred since they provide particularly excellent effects for activating the vinyl monomer and for firm consolidation with the polymer. It is one of the features of the present invention that the present invention is applicable to an inorganic compound to which usual coupling treatment is not applicable, and the shape and the size of the inorganic compound can suitably be selected.

As the vinyl monomer for the preparation of the organic composite filler (A) of the present invention, any usual radically polymerizable vinyl monomer may be used. Specifically, the vinyl monomer includes alkyl esters of (meth)acrylic acid such as methyl methacrylate and butyl acrylate; aromatic vinyl monomers such as styrene and α -methyl styrene; vinyl cyanide monomers such as acrylonitrile and methacrylonitrile; and monomers useful as the ethylenic vinyl monomer mixture (B) in the present invention. The monomer is sultably selected depending upon the particular purpose.

The radical polymerization initiator to be used for the preparation of the organic composite filler (A) may be any usual peroxide or azo compound. However, it is preferred to employ a radical polymerization initiator which is decomposable within a temperature range of from 40 to 100°C. Among them, peroxides such as benzoyl peroxide and potassium persulfate and azo compounds such as azobisisobutylonitrile are particularly effective and preferred in view of the graft rate of the resulting organic composite filler.

The concentration of the carboxylic acid monomer for the preparation of the organic composite filler (A) is usually from about 0.05 to 100% by weight, preferably from 0.1 to 50% by weight, more preferably from 0.5 to 30% by weight, based on the total weight of the inorganic compound and the vinyl monomer. In many cases, it is preferred to increase the amount of the carboxylic acid monomer as the vinyl monomer component increases. The weight ratio of the vinyl monomer or a mixture of vinyl monomers to the inorganic compound used may be varied within a wide range and is usually within a range of from about 500:1 to about 1:5, preferably from about 50:1 to about 1:1. When the polymerization is conducted in an aqueous medium, the amount of water is usually from about 1% to several hundreds times, preferably from about 10% to 10 times, based on the total weight of the inorganic compound and the vinyl monomer.

The reaction is preferably conducted in an atmosphere of an inert gas such as nitrogen usually at a temperature of from about 10 to 100°C, preferably at a temperature of about 40 to 100°C at which the radical polymerization initiator can be decomposed. Here, the practical reaction temperature is suitably selected depending upon the vinyl monomer. However, it is important to conduct the reaction at a controlled temperature where thermal polymerization is negligible. If the reaction is conducted at a high temperature where thermal polymerization takes place substantially, the consolidation and the uniformity of the resulting composite filler will be impaired. The reaction time is usually from 30 minutes to about 15 hours. The resulting organic composite filler may be dried usually within a temperature range of from about 10 to 300°C, preferably from about 50 to 200°C. The interaction between the surface of the inorganic compound and the polymer provided by the method of the present invention is beyond a usual physical adhesion due to simple adsorption or van der Waals force. This is evident from the fact that a substantial amount of non-extracted polymer remains even when the vinyl polymer is subjected to extraction treatment with a good solvent.

In the above-mentioned organic composite filler (A), various inorganic compounds can be firmly

consolidated with an organic polymer. Thus, the compatibility of the inorganic compound and the organic polymer in the composite resin is good, and the inorganic compound required for a dental composition can optionally be selected depending upon the particular purpose to obtain a dental composition as the desired product of the present invention.

The ethylenic vinyl monomer mixture (B) to be used in the present invention contains Bis-MEPP and a hexa-functional urethane (meth)acrylate of the formula III. By the combination of the two types of monomers, it is surprisingly possible to obtain a composite resin having excellent mechanical strength and good water resistance when the composition of the present invention is cured.

Among the monomers of the formula III, a hexa-functional urethane acrylate wherein n is 8, R₁ is a hydrogen atom and R₂ is a methyl group (hereinafter referred to simply as U-6HA) and a hexa-functional urethane methacrylate wherein n is 8 and each of R₁ and R₂ is a methyl group (hereinafter referred to simply as U-6H) are preferred from the viewpoint of curability. Among them, U-6HA is most preferred.

In the present invention, there is no particular restriction as to the proportions of Bis-MEPP, the hexafunctional urethane (meth)acrylate and the diluting ethylenic vinyl monomer in the monomer mixture (B). However, from the viewpoint of the operation efficiency for handling the composition of the present invention, it is preferred to adjust the proportions so that the viscosity of the composition of the present invention would be from 1,000 to 500,000 poise at 25°C. Usually, the proportions of Bis-MEPP and the hexa-functional urethane (meth)acrylate in the monomer mixture (B) are from 40 to 70% by weight and from 10 to 40% by weight, respectively. Further, from the viewpoint of the water resistance and mechanical strength, the blend ratio (weight ratio) of the hexa-functional urethane (meth)acrylate to Bis-MEPP is usually from 0.05:1 to 1:1, preferably from 0.1:1 to 0.8:1, more preferably from 0.2:1 to 0.5:1.

As the diluting ethylenic vinyl monomer for the present invention, the conventional monomers commonly employed for dental composite resins may be used. Specific examples include alkanediol di(meth)-acrylates such as 1,6-hexamethylene glycol dimethacrylate; polyalkylene glycol di(meth)acrylates such as triethylene glycol dimethacrylate (hereinafter referred to simply as 3G) and tetraethylene glycol di(meth)-acrylate; alkyl esters of (meth)acrylate acid such as trimethylolpropane tri(meth)acrylate, glycidyl (meth)-acrylate, methyl methacrylate and butyl acrylate; aromatic vinyl compounds such as styrene and a-methyl styrene; and vinyl cyanide compounds such as acrylonitrile and methacrylonitrile. In the present invention, the radical polymerizable vinyl monomer used in the organic composite filler (A) and the monomer mixture (B) may be the same.

The proportions of the organic composite filler (A) and the monomer mixture (B) may optionally be selected depending upon the types and the purposes of the inorganic compound used for the filler (A) and the monomer mixture (B). However, the organic composite filler (A) and the monomer mixture (B) are used usually within the ranges of from 20 to 90 parts by weight and from 10 to 80 parts by weight, respectively.

There is no particular restriction as to the photopolymerization initiator (C) in the present invention. However, in view of its dental use in the oral cavity, it is preferably operable with a visible light having a wavelength of from 400 to 800 nm excluding the near ultraviolet region. Further, from the clinical standpoint, it is preferably curable in about 30 seconds.

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From the foregoing points, preferred photopolymerization initiators (C) include a-diketone compounds such as camphor quinone (hereinafter referred to simply as CQ), benzyl and diacetyl, and a mixture thereof. The a-diketone compound can be used in combination with a reducing agent. In the composition of the present invention, the reducing agent is required from the viewpoint of high curability to be diaminobenzophenone compounds of the formula:

$$R_1R_2N \longrightarrow C \longrightarrow NR_3R_4 \qquad (IV)$$

wherein each of R_1 to R_4 which may be the same or different is a hydrogen atom or a C_1 - C_3 alkyl group, provided that at least one of them is the alkyl group. Among them, 4.4'-bis(dimethylamino)benzophenone (hereinafter referred to simply as AB) wherein each of R_1 to R_4 is a methyl group is particularly preferred from the viewpoint of curability.

The amount of the photopolymerization initiator (C) is suitably selected taking into consideration the curability, the storage stability and the color tone of the composition. However, in a combined system of the a-diketone compound with the diaminobenzophenone compound, the initiator is used usually in such an amount that the a-diketone compound is from 0.01 to 15% by weight, preferably from 0.05 to 5% by

weight, more preferably from 0.1 to 2% by weight, relative to the monomer mixture (B), and the diaminobenzophenone compound is within a range of from 0.01 to 15% by weight, relative to the monomer mixture (B).

In addition to the above components, a filler may be added to the composition of the present invention, as the case requires. Such a filler component includes elements of Groups I, II, III, IV and V, transition metals and their oxides, hydroxides, chlorides, sulfates, sulfites, carbonates, phosphates and silicates, and mixtures and complex salts thereof. Among them, silicon dioxide, quartz powder, aluminum oxide, barium sulfate, titanium oxide, talc, glass powder, glass beads, glass fibers, a glass filler containing a barium salt or a lead salt, silica gel, colloidal silica, carbon fibers, zirconium oxide, tin oxide and other ceramics powders are preferred. The above filler may be a non-treated filler, a filler having its surface treated with e.g. a silane coupling agent or an organic composite filler obtained by consolidating a filler with a polymer, followed pulverization.

The amount of the filler to be incorporated can be suitably changed depending upon the particular purpose of the photopolymerizable dental composition. However, the filler is usually incorporated in an amount within a range of from 0.1 to $10^3\%$ by weight, preferably from 0.5 to $5 \times 10^2\%$ by weight, more preferably from 1 to $10^2\%$ by weight, relative to the above-mentioned monomer mixture (B) to form a paste composition.

Further, a coloring agent, a polymerization inhibitor (such as hydroquinone, methoxybenzophenone, methylphenol or hydroquinone monomethyl ether), an oxidation stabilizer, an ultraviolet absorber (such as benzophenone), a pigment (such as iron oxide or titanium oxide) and a dyestuff may further be incorporated to the composition of the present invention, as the case requires.

Now, the present invention will be described in further detail with reference to Examples. However, it should be understood that the present invention is by no means restricted to such specific Examples. In the Examples, "parts" means "parts by weight".

Preparation of organic composite fillers

Organic composite filler (1):

In a 500 ml four necked separable flask equipped with a condenser, a nitrogen supply tube, a stirrer and a thermo couple for detecting the internal temperature, 50 g of fine quartz powder (prepared by pulverizing quartz A-2 manufactured by Tatsumori K.K. to an average particle size of from 1 to 2 µm) and 50 g of barlum glass (prepared by pulverizing barium glass #7724 manufactured by Coming Company to an average particle size of from 1 to 2 µm) were suspended and dispersed as the inorganic compounds in 300 ml of deionized water and flushed with nitrogen for 30 minutes. Then, 3.0 g of methyl methacrylate was added as the vinyl monomer while vigorously stirring under a nitrogen atomsphere. Then, the suspension was heated to 70 °C in a warm water bath. After confirming that the suspension was in a uniformly dispersed state, 0.5 g of acrylic acid as the carboxylic acid monomer was gradually added, and the polymerization reaction was conducted at the same temperature for 8 hours.

After the completion of the reaction, the product was collected by filtration under reduced pressure, thoroughly washed with delonized water and then dried by hot air drying at 100°C to remove the moisture to obtain about 100 g of an organic composite filler (1). The composite filler thus obtained had a polymer content of 2.5% as measured by a baking method, and when it was subjected to a Soxhlet extraction test for 50 hours by using hot benzene as the extracting solvent, the polymer content after the extraction was 2.0%. Thus, it was found that the majority of the polymer in the composite filler was extremely firmly consolidated to the surface of the fine quartz powder and barium glass. Separately, 100 g of each of the fine quartz powder and the barium glass was polymerized in the same manner independently, whereby the resulting product had the same polymer content as in the case of the combined system. Therefore, it is considered that the two inorganic fillers in the combined system have their surfaces uniformly consolidated with the polymer.

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Organic composite filler (2):

An organic composite filler (2) was prepared in the same manner as in the preparation of the organic composite filler (1) except that maleic anhydride was used instead of acrylic acid, and the evaluation was conducted in the same manner as in the case of the filler (1). The filler (2) thereby obtained showed the same results as the filler (1), thus indicating uniform consolidation.

Organic composite filler (3):

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Into a 500 ml four necked separable flask equipped with a condenser, a nitrogen supply tube, a stirrer and a thermo couple for detecting the internal temperature, 50 g of fine quartz powder (prepared by pulverizing quartz A-2 manufactured by Tatsumori K.K. to an average particle size of from 1 to 2 µm) and 50 g of barium glass (prepared by pulverizing barium glass #7724 manufactured by Corning Company to an average particle size of from 1 to 2 µm) were suspended and dispersed as the inorganic compounds in 300 ml of delonized water and flushed with nitrogen for 30 minutes. Then, 3.0 g of methyl methacrylate was added as the vinyl monomer while vigorously stirring under a nitrogen atomsphere. Then, the suspension was heated to 70 °C in a warm water bath. After confirming that the suspension was in a uniformly dispersed state, a solution prepared by dissolving 0.07 g of banzoyl peroxide as the radical polymerization initiator in 0.5 g of acrylic acid as the carboxylic acid monomer was gradually added, and the polymerization reaction was conducted at the same temperature for 8 hours.

After the completion of the reaction, the product was collected by filtration under reduced pressure, thoroughly washed with deionized water and then dried by hot air drying at 100°C to remove the moisture to obtain about 100 g of an organic composite filler (3). The composite filler thus obtained had a polymer content of 3.2% as measured by a baking method, and when it was subjected to a Soxhlet extraction test for 50 hours by using hot benzene as the extracting solvent, the polymer content after the extraction was as high as 3.0%. Thus, it was found that the majority of the polymer in the composite filler was extremely firmly consolidated to the surface of the fine quartz powder and barium glass. Separately, 100 g of each of the fine quartz powder and the barium glass was polymerized in the same manner independently, whereby the resulting product had the same polymer content as in the case of the combined system. Therefore, it is considered that the two inorganic fillers in the combined system have their surfaces uniformly consolidated with the polymer.

Organic composite fillers (4) to (6):

Organic composite fillers (4) to (6) were prepared in the same manner as the organic composite filler (3) except that maleic anhydride was used instead of acrylic acid, butyl acrylate was used instead of methyl methacrylate, and azobisisobutyronitrile was used instead of benzoyl peroxide, respectively, and the evaluation was conducted in the same manner as in the case of the filler (3). The fillers (4) to (5) thus obtained showed the same results as the filler (3), thus indicating that they were uniformly consolidated.

To 90 g of each of the organic composite fillers (1) to (6) obtained by the methods as described above, 10 g of Aerosil (trade mark) R-972 (hydrophobic amorphous silica manufactured by Deutsche Gold-und Silber-Scheideanstalt Vormals Roessler) was added as the viscosity regulating filler to obtain fillers (A) to (F).

EXAMPLE 1 to 6 and COMPARATIVE EXAMPLES 1 and 2

Each of various photopolymerizable dental compositions prepared to have the composition as identified in Table 1, was filled in a stainless steel mold having an inner diameter of 4 mm and a height of 6 mm, and a cover glass having a thickness of about 0.1 mm was fitted on each of the top and lower surfaces. Then, by using a visible light irradiator ("Optilux" manufactured by Minnesota Mining and Manufacturing Company) irradiation was conducted from both the top and lower surfaces for 30 seconds with a distance between the cover glass and the irradiator of 1 mm to obtain a cured product. The cured product thus obtained was stored in water at 37° C for 24 hours, whereupon the compression strength was measured. Further, the cured product was stored in water at 37° C for one week, whereupon the compression strength was evaluated as the water resistance strength. The strength was measured at a cross head speed of 1.0

mm/min. by using Tensilone (IS-500 manufactured by Shimadzu Corporation).

Further, a cured product obtained in the same manner by using a stainless steel mold having an inner diameter of 20 mm and a height of 1 mm was stored in water at 37 °C for one week. Then, the weight increase per unit surface area of the cured product was measured to evaluate the water absorption.

For the purpose of comparison, similar evaluation was conducted with respect to a case (a) wherein 10 g of Aerosii (trade mark) R-972 was mixed to 90 g of a silane-treated filler obtained by treating a filler mixture comprising 50 g of fine quartz powder and 50 g of barium glass with 3% of a silane coupling agent (3-methacryloxypropyltrimethoxysilane, KBM #503 manufactured by Shinetsu Silicon K.K.), instead of the composite filler of the present invention and a case of a non-treated filler (b) prepared by mixing 10 g of Aerosii (trade mark) R-972 to 45 g of each of non-treated fine quartz powder and barium glass.

It is evident from Table 1 that dental materials using the composite fillers (A) to (F) of the present invention (Examples 1 to 6) provide cured products having superior mechanical strength and water resistance as compared with those wherein the conventional silane-treated fillers (Comparative Example 1) or non-treated fillers (Comparative Example 2) are used.

Further, the composite fillers used in the present invention had extremely good wettability with the monomer mixtures (B) and could readily be consolidated with the mixtures (B) to form uniform composites. Besides, the outer appearance of the cured products thereby obtained were substantially superior to the Comparative Examples.

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EXAMPLE 7 and COMPARATIVE EXAMPLES 3 and 4

Aqueous heterogeneous polymerization was conducted in the same manner as in Example 1 except that 100 g of aluminum exide powder (special grade reagent) as the inorganic compound to obtain an organic composite filler (G). The composite filler thus obtained had a polymer content of 2.5%, and the polymer content after the extraction treatment was 2.0%. By using this organic composite filler (G), dental compositions were prepared to have the compositions as identified in Table 2, and various physical properties of the cured products were evaluated in the same manner as in Example 1. The results thereby obtained are shown in Table 2.

For the purpose of comparison, evaluation in the same manner as in Example 1 was conducted with respect to a case wherein a siliane treated filler (c) obtained by treating aluminum oxide powder with 3% of a siliane coupling agent was used instead of the composite filler of the present invention and a case wherein non-treated aluminum oxide powder (d) was used instead of the composite filler of the present invention.

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EXAMPLE 8

Aqueous heterogeneous polymerization was conducted in the same manner as in Example 3 except that 100 g of aluminum oxide powder (special grade reagent) was used as the inorganic compound to obtain an organic composite filler (H). The composite filler thus obtained had a polymer content of 3.2%, and the polymer content after the extraction treatment was 2.9%. By using this organic composite filler (H), dental compositions were prepared to have the compositions as identified in Table 2, and various physical properties of the cured products were evaluated in the same manner as in Example 3. The results thereby obtained are shown in Table 2.

It is evident from Table 2 that like the results in the Table 1, those wherein the dental compositions of the present invention were used have superior properties as compared with those wherein the conventional silane-treated fillers or non-treated fillers were employed.

EXAMPLES 9 to 11 and COMPARATIVE EXAMPLES 5 to 7

Dental compositions were prepared in the same manner as in Example 1 except that the composition of the monomer mixture (B) was changed as shown in Table 3, and various physical properties of the cured products thereby obtained were evaluated in the same manner as in Example 1. The results are shown in Tables 3.

EXAMPLES 12 TO 16 and COMPARATIVE EXAMPLES 8 TO 12

Dental compositions were prepared in the same manner as in Example 3 except that the compositions of the monomer mixture (B) was changed as shown in Table 3, and various physical properties of the cured products thereby obtained were evaluated in the same manner as in Example 3.

The results are shown in Table 3.

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It is evident from Table 3 that by using specific monomers as the monomer mixture (B), it is possible to obtain cured products for dental use having excellent mechanical strength and water resistance.

As described in detail in the foregoing, the composition of the present invention is excellent in the mechanical strength and water resistance and when used as a dental material composition, it exhibits excellent effects.

Table 1

		Сотр	Composition of photopolymerizableental composition	opolymer!	zabident	alcompositio	Ul	Physical	Physical propertiesof cured product	edproduct
	T	Filler(A)	Monomer mixture (B) (parts)	nixture(B)(parts)	Photopol: Initiator(C	Photopolymerization Initiator(C)(parts)*1)	Compressive strength (kg/cm²)	Water absorption (mg/om*)	Water resistance strangth (kg/cm²)
.,.	Туре	Amount (parts)	Bis-MEPP	И-6НА	36	ბე	AB			
Example 1	A	70	18	. 9	19	0.4	2.0	3310	0.18	3310
Example 2	ω	20	18	ဖ	ø	0.4	2.0	3300	0.17	3290
Example3	ပ	20	18	69	9	0.4	2.0	3380	0.20	3380
Example 4	Δ	20	18	9	60	0.4	2.0	3350	0.17	3340
Example 5	m	02	18	9	9	04	20	3350	0,19	3350
Example	Ľ.	20	18	9	9	0.4	2.0	3370	0.18	3360
*1)Amountrelativeto 100 parts of the m	stativeto	100 parts of	the monomer	onomermixture(B)						

Table 1 (continued)

		Сотр	Composition photopolymerizabidentel composition	polymeriz	abldent	alcompositio	ı	Physical	Physical propertiesof cured product	ed product
		Filler(A)	Monomer mixture(B) (parts)	kture(B)(parts)	Photopoly initiator(C	Photopolymerization initiator(C)(parts)*1)	Compressive strangth(kg/cm²)	Water absorption (mg/cm*)	Water resistance strength(kg/cm²)
	Туре	Type Amount (parts)	віз-мерр и-вна за	U-6HA	36	ĊЭ	АВ			
ComparativeExample 1	В	0,2	18	8	9	4.0	2.0	. 2920	0.44	2490
ComparativeExample 2	۵	92	48	9	8	0.4	2.0	2180	0.33	1980

*1) Amountrelativeto 100 parts of the monomer mixture (B).

Table 2

Type Amount Bis-MEPP U-6HA 3G (parts) G 30 42 14 14 14 C 30 42 14 14 14 C 30 42 14 14 14 14 14 14 14 14 14 14 14 14 14	Composition protopolymenzapieentarcomposition	Physical	Physical properties of cured product	edproduct
Type Amount Bis-MEPP U-6HA 3G April Apri	Photopolymerization Initiator(С)(ратз)*1)	Compressive strength(kg/cm²)	Water absorption (mg/cm²)	Water resistance strength(kg/cm²)
G 30 42 14 14 14 14 14 30 42 14 14 14 14 14 14 14 14 14 14 14 14 14				-
veExample 3 C 30 42 14 14 14 14 14 14 14		1040	0.21	1040
30 42 14 14	0.4 2.0	1070	0.24	1050
		380	0.58	280
45 46 14		420	0.45	300

Table 3

<u> </u>		Compositional		photopolymerizabidentalcomposition	bidentai	d L L L	osition		Fnysical	riiysicai pi opei ilesoi cui eu pi ouuti	anbloam:
	Filler(A)	×	Monomer mixt.	ıer mixtura(B) (parts)	arts)		Photopoly iniliator(C	Photopolymerization initiator(C)(parts)*1)	Compressive strength (kg/cm*	Water absorption (mg/cm³)	Water resistance strength(kg/cm²)
Туре	Amount (parts)	ВІЗ-МЕРР	BIS-GMA	U-6HA	п-вн	36	g	AB			
Example 9 A	2	18	•	•	8	8	0.4	2.0	3320	0.18	3320
Example 10 A	20	=	,	9	,	9	0.4	2,0	3210	0.32	3200
Example 11 A	20	22	,	2	•	.00	0.4	2.0	3010	0.19	3000
Example 12 C	2	18	,		0	9	0.4	2.0	3370	0.17	3360
Example 13 C	70	-	ı	.	,	9	0.4	2.0	3320	0.28	3310
	2	22	•	2	,	9	0.4	2.0	3100	0.18	3100

*1)Amountrelativeto 100 parts of the monomer mixture (B).

Table 3 (continued)

			Composition photopolymerizable antal composition	of photopo	lymeriza	bldent	alcon	position		Physicalp	Physical properties of cured product	edproduct
	Ē	Filler(A)	Mor	Monomermixture(B)(parts)	ıre(B)(p	arts)		Photopolymerizallor initiator(C)(parts)*1)	Photopolymerization initiator(C)(parts)*1)	Compressive strength (kg/cm²)	Water absorption (mg/cm²)	Water resistance strength (kg/cm²)
	Туре	Amount (parts)	Type Amount Bis-MEPP (parts)	BIS-GMA U-8HA U-6H 3G	U- ВНА	H9-D	36	8	AB			
Example 15	ပ	0%	R	1	4	٠	9	0.4		3360	0.19	3350
Example 18	ပ	2	16	,	1	80	ဖ	0.4	2.0	3370	0.17	3370
ComparativeExamples	«	2	24	1	,	·	9	0.4	5.0	2710	0.19	2670
ComparativeExample8	4	2	- 3-	24	,	•	9	0.4	2.0	3210	0.62	2860
ComparativeExample7	⋖	2	52	12		ı	9	0.4	2.0	2820	0.43	2480
*1) Amountrelativeto 100 parts of the monomer) parts	of the mor	nomermixture(B).	-a(B).								

Table 3 (continued)

	S	Composition of photopolymerizabidental composition	of photopol	ymerizal	oldenta	loom	position		Physical	Physical propertiesof cured product	redproduct
	Filler (A)	Mon	Monomer mixture (B) (parts)	re(6)(pa	rts)		Photopoly initiator(C)	Photopolymerization initiator(C)(parts)*1)	Compressive strength (kg/cm²)	Water absorption (mg/cm²)	Water resistance strength(kg/cm²)
Турв	Amount (parts)	Type Amount Bis-MEPP (parts)	BIS-GMA U-6HA U-6H 3G	U-6HA	H8-7	36	. 60	AB			
ComparativeExample8 C	02	24		•	•	8	0.4	2.0	2800	0.18	2800
ComparativeExample9 C	20	1	8	•	•	9	0.4	2.0	3330	0.47	3010
ComparativeExample 10 C	20	42	12	,	•	9	0.4	20	. 2950	0.42	2800
ComparativeExample 11 C	20	•	82	9	,	9	0.4	20	3380	0.46	3060
ComparativeExample 12 C	92		81	,	9	0	0.4	2.0	- 3330	0.47	3020

*1) Amount relative to parts of the monomer mixture (B),

Claims

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1. A photopolymerizable dental composition, which comprises:

(A) an organic composite filler obtained by polymerizing at least one carboxylic acid monomer of the formula:

$$R_2 = C = R_3$$

$$C = C COOX$$

wherein R₁ is a hydrogen atom, a C₁-C₁₅ alkyl group, a -COOY group, a halogen atom, or a phenyl group or a derivative thereof; R₂ is a hydrogen atom, a C₁-C₁₅ alkyl group, a -COOZ group, a halogen atom, or a phenyl group or a derivative thereof; R₃ is a hydrogen atom, a C₁-C₁₅ alkyl group, a halogen atom, or a phenyl group or a derivative thereof; and each of X, Y and Z is a hydrogen atom, NH₄ or an alkali metal atom; or the formula:

wherein each of R₄ and R₅ is a hydrogen atom, a C₁-C₁₅ alkyl group, a halogen atom, or a phenyl group or a derivative thereof; and at least one radical polymerizable vinyl monomer in a polymerization system in which an inorganic compound is dispersed;

(B) a monomer mixture comprising 2.2-bis[4-(methacryloxyethoxy)phenyl]propane, at least one hexafunctional urethane (meth)acrylate of the formula:

wherein X is
$$\{CH_2\}_{n}$$
 HNCOO-CH $\{CH_2\}_{n}$ CH $_2$ OOC-C=CH $_2$

wherein n is an integer of from 1 to 10, and each of R1 and R2 which may be the same or different is a

hydrogen atom or a methyl group, and a diluting ethylenic vinyl monomer; and

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(C) a photopolymerization initiator comprising an a-diketone compound and a reducing agent, wherein the reducing agent is a diaminobenzophenone compound of the formula:

$$R_1R_2N$$
 NR_3R_4 (IV.)

wherein each of R₁ to R₄ which may be the same or different is a hydrogen atom or a C₁-C₅ alkyl group, provided that at least one of them is the alkyl group.

- 2. The photopolymerizable dental composition according to Claim 1, wherein the carboxylic acid monomer is selected from the group consisting of acrylic acid, methacrylic acid, crotonic-acid and maleic anhydride.
- 3. The photopolymerizable dental composition according to Claim 1, wherein the inorganic compound is at least one member selected from the group consisting of barium sulfate, barium fluoride, silicon dioxide, aluminum oxide, titanium oxide, quartz powder, glass powder, glass beads, glass fibers, a glass filler containing a barium salt, a lead salt or a strontium salt, silica gel, zirconium oxide and tin oxide.
- 4. The photopolymerizable dental composition according to Claim 1, wherein the a-diketone compound is at least one member selected from the group consisting of camphor quinone, benzyl and diacetyl.

EUROPEAN SEARCH REPORT

		SIDERED TO BE RELEVANT		EP 88109469.2
Category	Citation of document w	ith indication, where appropriate, vant passages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. CI.4)
A	EP - A2/A3 - C CHEMICAL INDUS * Claims 1.	•	1,4	A 61 K 6/08
	•		•	
A	CHEMICAL INDUS	2 535 (IMPERIAL TRIES LIMITED)	1,4	
	* Claims 1,	7 *		
A	EP - A2 - 0 01	7 937 (KANEBO LTD.)	1	
A	EP - A2 - 0.15	 5 812 (KURARAY CO	1.4	
	LTD.)	8; page 20.	-, .	
				TECHNICAL FIELDS SEARCHED (Int. CI.4)
A	EP - A2/A3 - O HAAS COMPANY)	104 000 (ROHM AND	1	A 61 K
	* Claim 1 *			
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	The present search report has b	een drawn up for all claims		
	Place of search VIENNA	Date of completion of the search 14–09–1988		Exeminer HEIN
Y: parti	CATEGORY OF CITED DOCL cularly relevant if taken alone cularly relevant if combined waters of the same category relogical background written displaying	E: earlier pater	nciple under it document, ig date ted in the ap ted for other	rlying the invention
O: non-	written disclosure mediate document	& : member of to	he same pate	ent family, corresponding

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(1) Publication number:

0 295 627 B1

(P).

EUROPEAN PATENT SPECIFICATION

- Date of publication of patent specification: 30.12.92 (5) Int. Cl.5. A61K 6/08
- 21 Application number: 88109469.2
- 2 Date of filing: 14.06.88
- Photopolymerizable dental composition.
- Priority: 19.06.87 JP 153022/87
- ② Date of publication of application: 21.12.88 Bulletin 88/51
- 49 Publication of the grant of the patent: 30.12.92 Bulletin 92/53
- Designated Contracting States:
 DE FR GB
- 66 References cited:

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EP-A- 0 155 B12

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- Proprietor: MITSUBISHI RAYON CO., LTD. 3-19, Kyobashi 2-chome Chuo-Ku Tokyo 104(JP)
- inventor: Sasaki, Isao
 1665-98, Ohno-cho
 Saiki-gun Hiroshima-ken(JP)
 Inventor: Mukal, Nobuhiro
 13-19-105, Inokuchidai 1-chome
 Nishi-ku Hiroshima-shi Hiroshima-ken(JP)
 Inventor: Ige, Hitoshi
 3-16-303, Nishisakae 3-chome
 Ohtake-shi Hiroshima-ken(JP)
- Representative: TER MEER MÜLLER STEIN-MEISTER & PARTNER Mauerkircherstrasse 45 W-8000 München 80(DE)

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Description

The present invention relates to a dental filling composition containing a substantial amount of an organic composite filler as inorganic material. More particularly, it relates to a photopolymenizable dental filling composition which can readily be polymerized by irradiation with a visible light to provide a cured product having high mechanical properties and excellent water resistance.

For dental filling material, it is important that such material has not only good physical properties such as mechanical strength, abrasion resistance, water resistance and adhesive properties but also aesthetical properties such as transparency and polishability to make it look like natural teeth. Many attempts have been made to develop a so-called composite resin having an inorganic filler incorporated into a resin, as a substitute material for amalgam and with an object to improve the above-mentioned properties.

This composite resin is usually composed essentially of three components of (1) an inorganic filler, (2) an olefinic unsaturated compound and (3) a polymerization initiator. However, if an inorganic filler and an organic resin are simply blended, it is difficult to obtain adequate composite properties since the inorganic filler and the organic resin are poor in the interfacial affinity such as the compatibility or the adhesion to each other. In order to overcome such a drawback, a dental material has been proposed in which a filler prepared by treating a glass surface with a silane coupling agent is incorporated. However, the surface treatment with a silane coupling agent is limited in its applicable inorganic material to a glass filler having silanol groups on its surface and thus has a serious drawback that it is poor in the general applicability. Further, there is a problem such that the formed siloxane linkage is susceptible to hydrolysis and thus is poor in the water resistance. Accordingly, when applied to a dental filling composition, it hardly provides an interfacial reinforcing property to a quartz filler which is commonly employed as a filler to impart primarily high hardness and aesthetical properties. Further, it has a clinical problem such that since the water resistance is poor, the mechanical strength tends to deteriorate as time passes during clinic for a long period of time in a wet oral cavity.

Further, various studies have been made on the monomer constituting the organic resin. For example, a bisphenol A multi-functional monomer represented by bisphenol A diglycidyl methacrylate (hereinafter referred to simply as Bis-GMA) has been proposed as a monomer which provides a composite having a small polymerization shrinkage factor, and a hydrophobic monomer 2,2-bis[4-(methacryloxyethoxy)-phenyl]-propane (hereinafter referred to simply as Bis-MEPP) has been proposed as a monomer which provides excellent water resistance. However, Bis-GMA has a hydroxyl group in its molecule, and the resulting cured product is highly water absorptive and thus is poor in the water resistance. On the other hand, when Bis-MEPP is used, the curability is poor and adequate crosslinking density is hardly obtainable, and there has been a problem that the product is poor in the mechanical strength. Further, even if Bis-GMA and Bis-MEPP are combined to complement these drawbacks, it is impossible to obtain a composite resin having satisfactory properties. Accordingly, in order to determine a monomer composition for a particular purpose, it is necessary to study a very wide range of monomer compositions.

It is an object of the present invention to provide a photopolymerizable dental composition having excellent mechanical strength and good water resistance.

The present invention provides a photopolymerizable dental composition, which comprises;

(A) an organic composite filler obtained by polymerizing at least one carboxylic acid monomer of the formula:

$$R_{1} = C = C$$

$$C = C$$

$$C = C$$

$$C = C$$

$$C = C$$

wherein R_1 is a hydrogen atom, a C_1 - C_{15} alkyl group, a -COOY group, a halogen atom, or a phenyl group or a derivative thereof, R_2 is a hydrogen atom, a C_1 - C_{15} alkyl group, a -COOZ group, a halogen atom, or a phenyl group or a derivative thereof, R_3 is a hydrogen atom, a C_1 - C_{15} alkyl group, a halogen atom, or a phenyl group or a derivative thereof, and each of X, Y and Z is a hydrogen atom, NH, or an alkali metal atom; or the formula:

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wherein each of R₄ and R₅ is a hydrogen atom, a C₁-C₁₅ alkyl group, a halogen atom, or a phenyl group or a derivative thereof, and at least one radical polymerizable vinyl monomer in a polymerization system in which an inorganic compound is dispersed;

(B) a monomer mixture comprising 2,2-bis[4-(methacryloxyethoxy)phenyl]propane, at least one hexafunctional urethane (meth)acrylate of the formula:

wherein X is

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$$-(\text{CH}_2)_{\overline{\Pi}} \text{HNCOO-CH} \underbrace{\begin{array}{c} \text{CH}_2\text{OOC-C=CH}_2\\ \text{CH}_2\text{OOC-C=CH}_2\\ \text{R}_2 \end{array}}_{\text{R}_2}$$

wherein n is an integer of from 1 to 10, and each of R₁ and R₂ which may be the same or different is a hydrogen atom or a methyl group, and a diluting ethylenic vinylmonomer; and

(C) a photopolymerization initiator comprising an α -diketone compound and a reducing agent, wherein the reducing agent is a diaminobenzophenone compound of the formula:

$$R_1R_2N$$
 NR_0R_4 (IV)

wherein each of R₁ to R₄ which may be the same or different is a hydrogen atom or a C₁-C₅ alkyl group, provided that at least one of them is the alkyl group.

EP-A-0 284 991 which is comprised in the state of the art in the sense of Art. 54(3) EPC discloses a photopolymerizable dental composition similar to the composition of the present invention, except that in this earlier application dimethylaminoethylmethacrylate or alkylesters of 4-(N,N-dimethylamino) benzoic acid are used as reducing agents in the photopolymerization initiator.

Now, the present invention will be described in detail with reference to the preferred embodiments.

The organic composite filler (A) constituting the composition of the present invention can be obtained by polymerizing the carboxylic acid monomer with at least one other polymerizable vinyl monomer in a polymerization system in which an inorganic compound is dispersed. There is no particular restriction as to the method for its preparation. As a preferred embodiment, a method may be mentioned wherein a vinyl

monomer and an inorganic compound are suspended and dispersed in an aqueous medium under such a temperature condition that no thermal polymerization takes place, and then, the carboxylic acid monomer is added and stirred to initiate aqueous heterogeneous polymerization, whereby the polymerization is conducted for a predetermined period of time.

In the above polymerization, a radical polymerization initiator may be added as the case requires. For example, it is preferred to add a radical polymerization initiator when a high graft rate is required or when a mixture of various vinyl monomers is used.

The specific carboxylic acid monomer represented by the formula I or II to be used for the formation of the organic composite filler (A) is required to have a carboxylic acid group as an active site which brings about the polymerization activity and a double bond as an active site which provides firm consolidation of the resulting polymer and the inorganic compound. Any compound of the formula I or II having such functional groups can be used. Specifically, acrylic acid, methacrylic acid, crotonic acid, tiglic acid, cinnamic acid, maleic anhydride and citraconic anhydride may be mentioned. Among them, acrylic acid, methacrylic acid, crotonic acid and maleic anhydride are particularly preferred since they have very high polymerization activities.

The inorganic compound to be used for the preparation of the organic composite filler (A) includes elements of Groups I, II, III, IV and V of the Periodic Table, transition metals and their oxides, hydroxides, chlorides, sulfates, sulfates, carbonates, phosphates and silicates, and mixtures and complex salts thereof. Among them, barium sulfate, barium fluoride, silicon dioxide, alminum oxide, titanium oxide, quartz powder, glass powder, glass beads, glass fibers, a glass filler containing a barium salt, a lead salt or a strontium salt, silica gel, zirconium oxide and tin oxide are particularly preferred since they provide particularly excellent effects for activating the vinyl monomer and for firm consolidation with the polymer. It is one of the features of the present invention that the present invention is applicable to an inorganic compound to which usual coupling treatment is not applicable, and the shape and the size of the inorganic compound can suitably be selected.

As the vinyl monomer for the preparation of the organic composite filler (A) of the present invention, any usual radically polymerizable vinyl monomer may be used. Specifically, the vinyl monomer includes alkyl esters of (meth)acrylic acid such as methyl methacrylate and butyl acrylate; aromatic vinyl monomers such as styrene and α -methyl styrene; vinyl cyanide monomers such as acrylonitrile and methacrylonitrile; and monomers useful as the ethylenic vinyl monomer mixture (B) in the present invention. The monomer is suitably selected depending upon the particular purpose.

The radical polymerization initiator to be used for the preparation of the organic composite filler (A) may be any usual peroxide or azo compound. However, it is preferred to employ a radical polymerization initiator which is decomposable within a temperature range of from 40 to 100°C. Among them, peroxides such as benzoyl peroxide and potassium persulfate and azo compounds such as azobisisobutylonitrile are particularly effective and preferred in view of the graft rate of the resulting organic composite filler.

The concentration of the carboxylic acid monomer for the preparation of the organic composite filler (A) is usually from about 0.05 to 100% by weight, preferably from 0.1 to 50% by weight, more preferably from 0.5 to 30% by weight, based on the total weight of the inorganic compound and the vinyl monomer. In many cases, it is preferred to increase the amount of the carboxylic acid monomer as the vinyl monomer component increases. The weight ratio of the vinyl monomer or a mixture of vinyl monomers to the inorganic compound used may be varied within a wide range and is usually within a range of from about 500:1 to about 1:5, preferably from about 500:1 to about 1:1. When the polymerization is conducted in an aqueous medium, the amount of water is usually from about 1% to several hundreds times, preferably from about 10% to 10 times, based on the total weight of the inorganic compound and the vinyl monomer.

The reaction is preferably conducted in an atmosphere of an inert gas such as nitrogen usually at a temperature of from about 10 to 100°C, preferably at a temperature of about 40 to 100°C at which the radical_polymerization initiator_can be decomposed. Here, the practical reaction_temperature is suitably selected depending upon the vinyl monomer. However, it is important to conduct the reaction at a controlled temperature where thermal polymerization is negligible. If the reaction is conducted at a high temperature where thermal polymerization takes place substantially, the consolidation and the uniformity of the resulting composite filler will be impaired. The reaction time is usually from 30 minutes to about 15 hours. The resulting organic composite filler may be dried usually within a temperature range of from about 10 to 300°C, preferably from about 50 to 200°C. The interaction between the surface of the inorganic compound and the polymer provided by the method of the present invention is beyond a usual physical adhesion due to simple adsorption or van der Waals force. This is evident from the fact that a substantial amount of non-extracted polymer remains even when the vinyl polymer is subjected to extraction treatment with a good solvent.

In the above-mentioned organic composite filler (A), various inorganic compounds can be firmly consolidated with an organic polymer. Thus, the compatibility of the inorganic compound and the organic polymer in the composite resin is good, and the inorganic compound required for a dental composition can optionally be selected depending upon the particular purpose to obtain a dental composition as the desired product of the present invention.

The ethylenic vinyl monomer mixture (B) to be used in the present invention contains Bis-MEPP and a hexa-functional urethane (meth)acrylate of the formula III. By the combination of the two types of monomers, it is surprisingly possible to obtain a composite resin having excellent mechanical strength and good water resistance when the composition of the present invention is cured.

Among the monomers of the formula III, a hexa-functional urethane acrylate wherein n is 6, R₁ is a hydrogen atom and R₂ is a methyl group (hereinafter referred to simply as U-6HA) and a hexa-functional urethane methacrylate wherein n is 6 and each of R₁ and R₂ is a methyl group (hereinafter referred to simply as U-6H) are preferred from the viewpoint of curability. Among them, U-6HA is most preferred.

In the present invention, there is no particular restriction as to the proportions of Bis-MEPP, the hexafunctional urethane (meth)acrylate and the diluting ethylenic vinyl monomer in the monomer mixture (B).

However, from the viewpoint of the operation efficiency for handling the composition of the present invention, it is preferred to adjust the proportions so that the viscosity of the composition of the present invention would be from 100 to 50,000 Pa*s (1,000 to 500,000 poise) at 25°C. Usually, the proportions of Bis-MEPP and the hexa-functional urethane (meth)acrylate in the monomer mixture (B) are from 40 to 70% by weight and from 10 to 40% by weight, respectively. Further, from the viewpoint of the water resistance and mechanical strength, the blend ratio (weight ratio) of the hexa-functional urethane (meth)acrylate to Bis-MEPP is usually from 0.05:1 to 1:1, preferably from 0.1:1 to 0.8:1, more preferably from 0.2:1 to 0.5:1.

As the diluting ethylenic vinyl monomer for the present invention, the conventional monomers commonly employed for dental composite resins may be used. Specific examples include alkanediol di(meth)-acrylates such as 1,6-hexamethylene glycol dimethacrylate; polyalkylene glycol di(meth)acrylates such as triethylene glycol dimethacrylate (hereinafter referred to simply as 3G) and tetraethylene glycol di(meth)-acrylate; alkyl esters of (meth)acrylic acid such as trimethylolpropane tri(meth)acrylate, glycidyl (meth)-acrylate, methyl methacrylate and butyl acrylate; aromatic vinyl compounds such as styrene and α -methyl styrene; and vinyl cyanide compounds such as acrylonitrile and methacrylonitrile. In the present invention, the radical polymerizable vinyl monomer used in the organic composite filler (A) and the monomer mixture (B) may be the same.

The proportions of the organic composite filler (A) and the monomer mixture (B) may optionally be selected depending upon the types and the purposes of the inorganic compound used for the filler (A) and the monomer mixture (B). However, the organic composite filler (A) and the monomer mixture (B) are used usually within the ranges of from 20 to 90 parts by weight and from 10 to 80 parts by weight, respectively.

There is no particular restriction as to the photopolymerization initiator (C) in the present invention. However, in view of its dental use in the oral cavity, it is preferably operable with a visible light having a wavelength of from 400 to 800 nm excluding the near ultraviolet region. Further, from the clinical standpoint, it is preferably curable in about 30 seconds.

The photopolymerization initiators (C) include α -diketone compounds such as camphor quinone (hereinafter referred to simply as CQ), benzil and diacetyl, and a mixture thereof. The α -diketone compound is used in combination with a reducing agent. In the composition of the present invention, the reducing agent is required from the viewpoint of high curability to be diaminobenzophenone compounds of the formula:

$$R_1R_2N$$
 NR_3R_4 (IV)

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wherein each of R₁ to R₄ which may be the same or different is a hydrogen atom or a C₁-C₅ alkyl group, provided that at least one of them is the alkyl group. Among them, 4,4'-bis(dimethylamino)benzophenone (hereinafter referred to simply as AB) wherein each of R₁ to R₄ is a methyl group is particularly preferred from the viewpoint of curability.

The amount of the photopolymerization initiator (C) is suitably selected taking into consideration the curability, the storage stability and the color tone of the composition. However, in a combined system of the α -diketone compound with the diaminobenzophenone compound, the initiator is used usually in such an

amount that the α -diketone compound is from 0.01 to 15% by weight, preferably from 0.05 to 5% by weight, more preferably from 0.1 to 2% by weight, relative to the monomer mixture (B), and the diaminobenzophenone compound is within a range of from 0.01 to 15% by weight, relative to the monomer mixture (B).

In addition to the above components, a further filler may be added to the composition of the present invention, as the case requires. Such a filler component includes elements of Groups I, II, III, IV and V, transition metals and their oxides, hydroxides, chlorides, sulfates, sulfates, carbonates, phosphates and silicates, and mixtures and complex salts thereof. Among them, silicon dioxide, quartz powder, aluminum oxide, barium sulfate, titanium oxide, talc, glass powder, glass beads, glass fibers, a glass filler containing a barium salt or a lead salt, silica gel, colloidal silica, carbon fibers, zirconium oxide, tin oxide and other ceramics powders are preferred. The above filler may be a non-treated filler, a filler having its surface treated with e.g. a silane coupling agent or an organic composite filler obtained by consolidating a filler with a polymer, followed pulverization.

The amount of the further filler to be optionally incorporated can be suitably changed depending upon the particular purpose of the photopolymerizable dental composition. However, the filler is usually incorporated in an amount within a range of from 0.1 to 10³% by weight, preferably from 0.5 to 5 x 10²% by weight, more preferably from 1 to 10²% by weight, relative to the above-mentioned monomer mixture (B) to form a paste composition.

Further, a coloring agent, a polymerization inhibitor (such as hydroquinone, methoxybenzophenone, methylphenol or hydroquinone monomethyl ether), an oxidation stabilizer, an ultraviolet absorber (such as benzophenone), a pigment (such as iron oxide or titanium oxide) and a dyestuff may further be incorporated to the composition of the present invention, as the case requires.

Now, the present invention will be described in further detail with reference to Examples. However, it should be understood that the present invention is by no means restricted to such specific Examples. In the Examples, "parts" means "parts by weight".

Preparation of organic composite fillers

Organic composite filler (1):

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In a 500 ml four necked separable flask equipped with a condenser, a nitrogen supply tube, a stirrer and a thermo couple for detecting the internal temperature, 50 g of fine quartz powder (prepared by pulverizing quartz A-2 manufactured by Tatsumori K.K. to an average particle size of from 1 to 2 µm) and 50 g of barium glass (prepared by pulverizing barium glass #7724 manufactured by Corning Company to an average particle size of from 1 to 2 µm) were suspended and dispersed as the inorganic compounds in 300 ml of deionized water and flushed with nitrogen for 30 minutes. Then, 3.0 g of methyl methacrylate was added as the vinyl monomer while vigorously stirring under a nitrogen atomsphere. Then, the suspension was heated to 70 °C in a warm water bath. After confirming that the suspension was in a uniformly dispersed state, 0.5 g of acrylic acid as the carboxylic acid monomer was gradually added, and the polymerization reaction was conducted at the same temperature for 8 hours.

After the completion of the reaction, the product was collected by filtration under reduced pressure, thoroughly washed with deionized water and then dried by hot air drying at 100°C to remove the moisture to obtain about 100 g of an organic composite filler (1). The composite filler thus obtained had a polymer content of 2.5% as measured by a baking method, and when it was subjected to a Soxhlet extraction test for 50 hours by using hot benzene as the extracting solvent, the polymer content after the extraction was 2.0%. Thus, it was found that the majority of the polymer in the composite filler was extremely firmly consolidated to the surface of the fine quartz powder and barium glass. Separately, 100 g of each of the fine quartz powder and the barium glass was polymerized in the same manner independently, whereby the resulting product had the same polymer content as in the case of the combined system. Therefore, it is considered that the two inorganic fillers in the combined system have their surfaces uniformly consolidated with the polymer.

Organic composite filler (2):

An organic composite filler (2) was prepared in the same manner as in the preparation of the organic composite filler (1) except that maleic anhydride was used instead of acrylic acid, and the evaluation was conducted in the same manner as in the case of the filler (1). The filler (2) thereby obtained showed the same results as the filler (1), thus indicating uniform consolidation.

Organic composite filler (3):

Into a 500 ml four necked separable flask equipped with a condenser, a nitrogen supply tube, a stirrer and a thermo couple for detecting the internal temperature, 50 g of fine quartz powder (prepared by pulverizing quartz A-2 manufactured by Tatsumori K.K. to an average particle size of from 1 to 2 µm) and 50 g of barium glass (prepared by pulverizing barium glass #7724 manufactured by Corning Company to an "average particle size of from 1 to 2 µm) were suspended and dispersed as the inorganic compounds in 300 ml of deionized water and flushed with nitrogen for 30 minutes. Then, 3.0 g of methyl methacrylate was added as the vinyl monomer while vigorously stirring under a nitrogen atomsphere. Then, the suspension was heated to 70°C in a warm water bath. After confirming that the suspension was in a uniformly dispersed state, a solution prepared by dissolving 0.07 g of benzoyl peroxide as the radical polymerization initiator in 0.5 g of acrylic acid as the carboxylic acid monomer was gradually added, and the polymerization reaction was conducted at the same temperature for 8 hours.

After the completion of the reaction, the product was collected by filtration under reduced pressure, thoroughly washed with deionized water and then dried by hot air drying at 100°C to remove the moisture to obtain about 100 g of an organic composite filler (3). The composite filler thus obtained had a polymer content of 3.2% as measured by a baking method, and when it was subjected to a Soxhlet extraction test for 50 hours by using hot benzene as the extracting solvent, the polymer content after the extraction was as high as 3.0%. Thus, it was found that the majority of the polymer in the composite filler was extremely firmly consolidated to the surface of the fine quartz powder and barium glass. Separately, 100 g of each of the fine quartz powder and the barium glass was polymerized in the same manner independently, whereby the resulting product had the same polymer content as in the case of the combined system. Therefore, it is considered that the two inorganic fillers in the combined system have their surfaces uniformly consolidated with the polymer.

Organic composite fillers (4) to (6):

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Organic composite fillers (4) to (6) were prepared in the same manner as the organic composite filler (3) except that maleic anhydride was used instead of acrylic acid, butyl acrylate was used instead of methyl methacrylate, and azobisisobutyronitrile was used instead of benzoyl peroxide, respectively, and the evaluation was conducted in the same manner as in the case of the filler (3). The fillers (4) to (6) thus obtained showed the same results as the filler (3), thus indicating that they were uniformly consolidated.

To 90 g of each of the organic composite fillers (1) to (6) obtained by the methods as described above, 10 g of Aerosil (trade mark) R-972 (hydrophobic amorphous silica manufactured by Deutsche Gold-und Silber-Scheideanstalt Vormals Roessler) was added as the viscosity regulating filler to obtain fillers (A) to (F).

EXAMPLE 1 to 6 and COMPARATIVE EXAMPLES 1 and 2

Each of various photopolymerizable dental compositions prepared to have the composition as identified in Table 1, was filled in a stainless steel mold having an inner diameter of 4 mm and a height of 6 mm, and a cover glass having a thickness of about 0.1 mm was fitted on each of the top and lower surfaces. Then, by using a visible light irradiator ("Optilux" manufactured by Minnesota Mining and Manufacturing Company) irradiation was conducted from both the top and lower surfaces for 30 seconds with a distance between the cover glass and the irradiator of 1 mm to obtain a cured product. The cured product thus obtained was stored in water at 37 °C for 24 hours, whereupon the compression strength was measured. Further, the cured product was stored in water at 37 °C for one week, whereupon the compression strength was evaluated as the water resistance strength. The strength was measured at a cross head speed of 1.0 mm/min, by using Tensilone (IS-500 manufactured by Shimadzu Corporation).

Further, a cured product obtained in the same manner by using a stainless steel mold having an inner diameter of 20 mm and a height of 1 mm was stored in water at 37°C for one week. Then, the weight increase per unit surface area of the cured product was measured to evaluate the water absorption.

For the purpose of comparison, similar evaluation was conducted with respect to a case (a) wherein 10 g of Aerosil (trade mark) R-972 was mixed to 90 g of a silane-treated filler obtained by treating a filler mixture comprising 50 g of fine quartz powder and 50 g of barium glass with 3% of a silane coupling agent (3-methacryloxypropyltrimethoxysilane, KBM #503 manufactured by Shinetsu Silicon K.K.), instead of the composite filler of the present invention and a case of a non-treated filler (b) prepared by mixing 10 g of Aerosil (trade mark) R-972 to 45 g of each of non-treated fine quartz powder and barium glass.

It is evident from Table 1 that dental materials using the composite fillers (A) to (F) of the present invention (Examples 1 to 6) provide cured products having superior mechanical strength and water resistance as compared with those wherein the conventional silane-treated fillers (Comparative Example 1) or non-treated fillers (Comparative Example 2) are used.

Further, the composite fillers used in the present invention had extremely good wettability with the monomer mixtures (B) and could readily be consolidated with the mixtures (B) to form uniform composites. Besides, the outer appearance of the cured products thereby obtained were substantially superior to the Comparative Examples.

10 EXAMPLE 7 and COMPARATIVE EXAMPLES 3 and 4

Aqueous heterogeneous polymerization was conducted in the same manner as in Example 1 except that 100 g of aluminum oxide powder (special grade reagent) as the inorganic compound to obtain an organic composite filler (G). The composite filler thus obtained had a polymer content of 2.5%, and the polymer content after the extraction treatment was 2.0%. By using this organic composite filler (G), dental compositions were prepared to have the compositions as identified in Table 2, and various physical properties of the cured products were evaluated in the same manner as in Example 1. The results thereby obtained are shown in Table 2.

For the purpose of comparison, evaluation in the same manner as in Example 1 was conducted with respect to a case wherein a silane treated filler (c) obtained by treating aluminum oxide powder with 3% of a silane coupling agent was used instead of the composite filler of the present invention and a case wherein non-treated aluminum oxide powder (d) was used instead of the composite filler of the present invention.

EXAMPLE 8

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Aqueous heterogeneous polymerization was conducted in the same manner as in Example 3 except that 100 g of aluminum oxide powder (special grade reagent) was used as the inorganic compound to obtain an organic composite filler (H). The composite filler thus obtained had a polymer content of 3.2%, and the polymer content after the extraction treatment was 2.9%. By using this organic composite filler (H), dental compositions were prepared to have the compositions as identified in Table 2, and various physical properties of the cured products were evaluated in the same manner as in Example 3. The results thereby obtained are shown in Table 2.

It is evident from Table 2 that like the results in the Table 1, those wherein the dental compositions of the present invention were used have superior properties as compared with those wherein the conventional silane-treated fillers or non-treated fillers were employed.

EXAMPLES 9 to 11 and COMPARATIVE EXAMPLES 5 to 7

Dental compositions were prepared in the same manner as in Example 1 except that the composition of the monomer mixture (B) was changed as shown in Table 3, and various physical properties of the cured products thereby obtained were evaluated in the same manner as in Example 1. The results are shown in Tables 3.

EXAMPLES 12 to 16 and COMPARATIVE EXAMPLES 8 to 12

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Dental compositions were prepared in the same manner as in Example 3 except that the compositions of the monomer mixture (B) was changed as shown in Table 3, and various physical properties of the cured products thereby obtained were evaluated in the same manner as in Example 3.

The results are shown in Table 3.

It is evident from Table 3 that by using specific monomers as the monomer mixture (B), it is possible to obtain cured products for dental use having excellent mechanical strength and water resistance.

As described in detail in the foregoing, the composition of the present invention is excellent in the mechanical strength and water resistance and when used as a dental material composition, it exhibits excellent effects.

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Table 1

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	Compos		photo	ројушег	izab	le dental	tion of photopolymerizable dental composition	Physical properties of cured product	perties of c	ured product
· · · · · · · · · · · · · · · · · · ·	Fille	ler (A)	Mon x l E eq)	Monomer mixture (B) (parls)		Photopolymeri- zation initiat (parts) *1)	Photopolymeri- zation initiator (C) (parts) *1)	Compressive atrength {kg/cm ² }	Water absorption' (mg/cm ²)	Water resistance strength
	Туре	Amount (parts)	Bis- MEPP	U-611A	36	ð	AB	N/mm ²		(kg/cm²) N/mm²
Example 1	4	7.0	18	9	9	0.4	2.0	(3310)	0.18	(3310)
Example 2	В	•		E,		•		(3300)	0.17	(3290) 322.4
Example 3	၁		ı	=	2	•	e	(3380)	0.20	(3380) 337.2
Example 4	G	e	*	2			£	(3350)	0.17	(3340)
Example 5	(L)	•		<u>.</u> .	z	2		(1350) 328.3	0.19	(3350) 326.3
Example 6	Ŀ	ŧ		1				(3370)	0.18	(3360)

*1) Amount relative to 100 parts of the monomer mixture (B).

Table 1 (continued)

	Compos	-	photo	polymer	izab	le dental	composition	tion of photopolymerizable dental composition Physical properties of cured product	perties of c	ured product
	Fille	ler (A)	Mon. He is	Monomer mixture (D) (parts)		Photopolymeri- zation initiati (parts) *1)	Photopolymeri- zation initiator (C) (parts) *1)	Comproseive strength (kg/cm²)	Water absorption (mg/cm ²)	Water resistance atrength
	Туре	Amount (parts)	Bls- MEPP	BIS- U-6HA	36	8	AB			(kg/cm²) N/mw²
Comparative Example 1	85	07	18	٠ و٠	Q	0.4	2.0	(2920) 2 <i>86</i> .2	0.44	(2490) 2440
Comparative Example 2	م		£		8	:	E	(2180) 273.6	0.33	(1980) 734.0

41) Amount relative to 100 parts of the monomer mixture (8).

Table 2

	Сопро	sition of	photo	ројушег	izab	le dental	Composition of photopolymerizable dental composition	Physical proparties of cured product	parties of c	ured product
	i i	ler (A)	Mon xim eq)	Monomer mixture (B) (parts)	_	Photopolymeri- zation initiati (parts) *1)	or (C)	Compressive strength {kg/cm ² }	Water absorption' (mg/cm ²)	
	Type	Amount (parts)	Bia- Mepp	U-611A	35	Ê	AB	N/mm²		(kg/cm ⁻) N/mm ²
Example 7	0	30	42	14	14	0.4	2.0	(1040)	0.21	(1040)
Example 8	=		8	1.	z	B	•	(1070)	0.24	(1050)
Comparative Example 3	Q	8		8				(380) 37.2	0.58	(280) 27.4
Comparative Example 4	פ	8		E		ı		(420) 44.2	0.45	(300)

ullet1] Amount relative to 100 parts of the monomer mixture (B).

Table 3

	Compos	sition of photopolymerizable dental composition	photo	ројуше	rízab	le de	ntal	сошров	ition	Physical pro	Physical properties of cured product	ured produc
-	FILI	ler (A)	Mon (pa	Monomer mixture (B) (parts)	ixtur	e (B)		Photopoly- merization initiator	Photopoly- merization initiator (C)	Compressive strength	Water absorption	Water resistance strength
	Туре	Amount (parts)	Bis- NEPP	Bis- GMA	U- 611A	-1.59	S S	CQ AI	AB	(kg/cm ⁻) N/mm ²	(mg/cm)	(kg/ 2m ²) N/mm ²
Example 9	<	7.0	18	ı		9	9	0.4	2.0	(3320)	0.18	(3320)
Example 10	1	9	11	ነ	13	1	2	1		(3210)	0.32	(3200)
Example 11	z	ε	22	4	2	1		· #	•	(3010)	0.19	(3000)
Example 12	U	3	13	1		υ	2		E	(1370)	0.17	(3360)
Example 13	•	,	=		2	'	8	2	ı	(3320)	0.28	(3310) 324.4
Example 14		3	22	ı	7	1	•	•		(3100)	91.0	(3100)

*1) Amount relative to 100 parts of the monomer mixture (8).

Table 3 (continued)

	Compos	sition of photopolymerizable dental composition	photo	polyme	rizab	le dei	ntal	compos	Ition	Physical pro	Physical properties of cured product	ured produci
	Ē.	Filler (A)	Mon (pa	Monomer mixture (B) (parts)	Ixtur	e (n)		Photopoly-merization initiator (parts)	Photopoly- merization initiator (C) (parts) *!)	Compressive strength {kg/cm ² }	Water absorption (mq/cm ²)	Water resistance atrength
-	Туре	Amount (parts)	Bis- MEPP	Bis- CMA	U- 611A	-h 159	32	O O	AB	N/mm²		(kg/cm^2) N/mm^2
Example 15	U ·	7.0	20	1	4		9	0.4	2.0	(3360)	0.19	(3350)
Example 16	=	ð	16	',	1	60	я.	a		(3370)	0.17	(3370)
Comparative. Example 5	V	e .	. 24	t	•	ı	8	8	=	(2710) 265.6	0.19	(2670) 7-132
Comparative Example 6	*	в .	•	24	ı	1	a	2	2	(3210)	0.52	(2850)
Comparative Example 7	•	•	12	12	1	1	8			(2020) 276.4	0.43	(2480)

Table 3 (continued)

	Сошрс	Composition of photopolymerizable dental composition	photo	polyme	rizab	le de	ntal	сошрав	ition	Physical pro	perties of c	Physical properties of cured product
	FII	ller (A)	Mon (pa	Monomer mixture (A) (parts)	ixtur	a (D)		Photopoly- merization initiator	Photopoly- merization initiator (C)	Compressive strength {ka/cm ² }	Water absorption (mo/cm ²)	Water resistance strength
	Туре	Amount (parts)	nis- Mepp	Bis- CMA	U- 611A	119	ä	8	AB	N/mm²		(kg/cm²) ///mm²
Comparative Example 8	ບ	70	24	ı	1	,	φ .	0.4	2.0	(2800) 274.4	0.18	(2800)
Comparative Example 9	1	τ	a	24	r	1	8	\$		(324.4	0.47	(30107)
Comparative Example 10	2	- \$	12	12	ı	1	=			(2950)	0.42	(2800)
Comperative Example 11	2	3	ı	2	w ·	i	2	2	t	(3360) 324.3	0.45	(3060)
Comparative Example 12	8	i	ı	91	1 :	ø	:	1	*	(3330)	0.47	(3020)

*1) Amount relative to 100 parts of the monomer mixture (B).

55 Claims

A photopolymerizable dental composition, which comprises:
 (A) an organic composite filler obtained by polymerizing at least one carboxylic acid monomer of the

formula:

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wherein R_1 is a hydrogen atom, a C_1 - C_{15} alkyl group, a -COOY group, a halogen atom, or a phenyl group or a derivative thereof; R_2 is a hydrogen atom, a C_1 - C_{15} alkyl group, a -COOZ group, a halogen atom, or a phenyl group or a derivative thereof; R_3 is a hydrogen atom, a C_1 - C_{15} alkyl group, a halogen atom, or a phenyl group or a derivative thereof; and each of X, Y and Z is a hydrogen atom, NH₄ or an alkali metal atom; or the formula:

wherein each of R_4 and R_5 is a hydrogen atom, a C_1 - C_{15} alkyl group, a halogen atom, or a phenyl group or a derivative thereof; and at least one radical polymerizable vinyl monomer in a polymerization system in which an inorganic compound is dispersed;

(B) a monomer mixture comprising 2,2-bis[4-(methacryloxyethoxy)phenyl]propane, at least one hexafunctional urethane (meth)acrylate of the formula:

wherein X is

wherein n is an integer of from 1 to 10, and each of R₁ and R₂ which may be the same or different is a hydrogen atom or a methyl group, and a diluting ethylenic vinyl monomer; and (C) a photopolymerization initiator comprising an α-diketone compound and a reducing agent, wherein the reducing agent is a diaminobenzophenone compound of the formula:

$$R_1R_2N$$
 NR_3R_4 (IV)

wherein each of R_1 to R_4 which may be the same or different is a hydrogen atom or a C_1 - C_5 alkyl group, provided that at least one of them is the alkyl group.

- 70 2. The photopolymerizable dental composition according to Claim 1, wherein the carboxylic acid monomer is selected from the group consisting of acrylic acid, methacrylic acid, crotonic-acid and maleic anhydride.
- 3. The photopolymerizable dental composition according to Claim 1, wherein the inorganic compound is at least one member selected from the group consisting of barium sulfate, barium fluoride, silicon dioxide, aluminum oxide, titanium oxide; quartz powder, glass powder, glass beads, glass fibers, a glass filler containing a barium salt, a lead salt or a strontium salt, silica gel, zirconium oxide and tin oxide.
- 4. The photopolymerizable dental composition according to Claim 1, wherein the α-diketone compound is at least one member selected from the group consisting of camphor quinone, benzil and diacetyl.

Patentansprüche

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25 1. Photopolymerisierbare Dentalzusammensetzung, umfassend:

(A) einen organischen zusammengesetzten Füllstoff, welcher durch Polymerisieren mindestens eines Carbonsäuremonomeren der Formet:

$$R_2 = C = R_3$$

$$R_1 = C = C$$

$$COOX$$
(1)

worin R₁ ein Wasserstoffatom, eine C₁-C₁₅-Alkylgruppe, eine-COOY-Gruppe, ein Halogenatom oder eine Phenylgruppe oder ein Derivat hiervon; R₂ ein Wasserstoffatom, eine C₁-C₁₅-Alkylgruppe, eine-COOZ-Gruppe, ein Halogenatom oder eine Phenylgruppe oder ein Derivat hiervon; R₃ ein Wasserstoffatom, eine C₁-C₁₅-Alkylgruppe, ein Halogenatom oder eine Phenylgruppe oder ein Derivat hiervon; und X, Y und Z jeweils ein Wasserstoffatom, NH₄ oder ein Alkalimetallatom bedeuten; oder der Formel:

worin R₄ und R₅ jeweils ein Wasserstoffatom, eine C₁-C₁₅-Alkylgruppe, ein Halogenatom oder eine Phenylgruppe oder ein Derivat hiervon bedeuten; sowie mindestens eines radikalisch polymerisierbaren Vinylmonomeren in einem Polymerisationssystem, in welchem eine anorganische Verbindung dispergiert ist, erhalten worden ist;

(B) eine Monomermischung, umfassend 2,2-Bis[4-(methacryloxyethoxy)-phenyl]-propan, mindestens

ein hexafunktionelles Urethan(meth)acrylat der Formel:

worin X

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bedeutet,

worin n eine ganze Zahl von 1 bis 10 und R₁ und R₂, die gleich oder verschieden sein können, jeweils ein Wasserstoffatom oder eine Methylgruppe darstellen, und ein ethylenisches Vinylmonomer zur Verdünnung; und

(C) einen Photopolymerisationsinitiator, umfassend eine α-Diketonverbindung und ein Reduktionsmittel, wobei das Reduktionsmittel eine Diaminobenzophenonverbindung der Formel:

$$R_1R_2N \longrightarrow C \longrightarrow NR_3R_4 \qquad (IV)$$

ist, worin R₁ bis R₄, die gleich oder verschieden sein können, jeweils ein Wasserstoffatom oder eine C₁-C₅-Alkylgruppe bedeuten, mit der Maßgabe, daß mindestens eine dieser Gruppen die Alkylgruppe darstellt.

- Photopolymerisierbare Dentalzusammensetzung nach Anspruch 1, wobei das Carbonsäuremonomer aus der aus Acrylsäure, Methacrylsäure, Crotonsäure und Maleinsäureanhydrid bestehenden Gruppe ausgewählt ist.
- S. Photopolymensierbare Dentalzusammensetzung nach Anspruch 1, wobei die anorganische Verbindung mindestens ein Vertreter ist, welcher aus der aus Bariumsulfat, Bariumfluorid, Siliciumdioxid, Aluminiumoxid, Titanoxid, Quarzpulver, Glaspulver, Glaskügelchen, Glasfasern, einem ein Bariumsalz, Bleisalz oder Strontiumsalz enthaltenden Glasfüllstoff, Silicagel, Zirkonoxid oder Zinnoxid bestehenden Gruppe ausgewählt ist.
- Photopolymerisierbare Dentalzusammensetzung nach Anspruch 1, wobei die anDiketonverbindung mindestens ein Vertreter ist, welcher aus der aus Campherchinon, Benzil und Diacetyl bestehenden Gruppe ausgewählt ist.

Revendications

1. Composition dentaire photopolymérisable, qui comprend :

(A) une charge composite organique obtenue par polymérisation d'au moins un acide carboxylique monomère représenté par la formule :

 $R_2 = C = C$ Coox(I)

dans laquelle :

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- R₁ représente un atome d'hydrogène, un groupe alkyle en C₁-C₁₅, un groupe -COOY, un atome d'halogène, ou un groupe phényle ou un dérivé de celui-ci ;
- R₂ représente un atome d'hydrogène, un groupe alkyle en C₁-C₁₅, un groupe -COOZ, un atome d'halogène, ou un groupe phényle ou un dérivé de celui-ci ;
- R₃ représente un atome d'hydrogène, un groupe alkyle en C₁-C₁₅, un atome d'halogène, ou un groupe phényle ou un dérivé de celui-ci; et
- X, Y et Z représentent chacun un atome d'hydrogène, NH, ou un atome de métal alcalin ; ou représenté par la formule ;

dans laquelle R_k et R_5 représentent chacun un atome d'hydrogène, un groupe alkyle en C_{15} , un atome d'halogène, ou un groupe phényle ou un dérivé de celui-ci ;

et d'au moins un monomère vinylique polymérisable par voie radicalaire, dans un système de polymérisation dans lequel un composé minéral est dispersé;

(B) un mélange de monomères comprenant le bis[(méthacryloxyéthoxy)-4 phényl]-2,2 propane, au moins un uréthanne (méth)acrylate hexafonctionnel, représenté par la formule :

$$\begin{array}{c|c}
x & 0 \\
y & C \\
y & C \\
y & C
\end{array}$$
(III)

dans laquelle :

- X représente

$$\begin{array}{c} & & \\ & \downarrow 1 \\ + \text{CH}_2 \\ & \uparrow n \end{array} \\ \text{HNCOO-CH} \begin{array}{c} \text{CH}_2 \\ \text{CH}_2 \\ \text{OOC-C=CH}_2 \\ & \downarrow \\ & \text{R}_2 \end{array}$$

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où n est un entier de 1 à 10, et R₁ et R₂, qui peuvent être identiques ou différents, représentent chacun un atome d'hydrogène ou un groupe méthyle ;

et un monomère vinylique éthylénique de dilution ; et

(C) un initiateur de photopolymérisation comprenant un composé de type α-dicétone et un agent réducteur, dans lequel l'agent réducteur est un composé de type diaminobenzophénone représenté par la formule :

$$R_1R_2N \longrightarrow C \longrightarrow NR_3R_4 \qquad (IV)$$

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dans laquelle R_1 à R_4 , qui peuvent être identiques ou différents, représentent chacun un atome d'hydrogène ou un groupe alkyle en C_1 - C_5 , à la condition qu'au moins l'un d'eux soit le groupe alkyle.

- Composition dentaire photopolymérisable selon la revendication 1, dans laquelle l'acide carboxylique monomère est choisi dans le groupe constitué par l'acide acrylique, l'acide méthacrylique, l'acide crotonique et l'anhydride maléique.
- 3. Composition dentaire photopolymérisable selon la revendication 1, dans laquelle le composé minéral est au moins un élément choisi dans le groupe constitué par le sulfate de baryum, le fluorure de baryum, le dioxyde de silicium, l'oxyde d'aluminium, l'oxyde de titane, la poudre de quarts, la poudre de verre, les perles de verre, les fibres de verre, une charge de verre contenant un sel de baryum, un sel de plomb ou un sel de strontium, le gel de silice, l'oxyde de zirconium et l'oxyde d'étain.
- 4. Composition dentaire photopolymérisable selon la revendication 1, dans laquelle le composé de type dicétone est au moins un élément choisi dans le groupe constitué par la camphre quinone, le benzile et le diacétyle.

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